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(54) Copper salts for laser marking of thermoplastic compositions

(57) A laser markable thermoplastic composition which contains a copper phosphate salt such as copper phosphate, copper sulfate, cupric hydroxide phosphate and copper thiocyanate. The copper phosphate salt is preferably added in an amount ranging from about 0.1 to 5 parts by weight and the copper phosphate salt preferably has a particle size less than 10 μ m. The laser markable thermoplastic composition can be laser marked to provide a visibly distinct and separately identifiable region which preferably differs in overall color from the base material by a Delta E value of at least 10-20.

low energy laser activation
of Cu salt to cause colour
change. Cu salt itself gives
colour. Optional additional
colorants but these absorb
light @ same @ Cu &
no disclosure of them causing
marking.

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Description**BACKGROUND OF THE INVENTION**

5 It is known to make a polymeric material which can be inscribed by means of laser light by mixing in a filler which changes color under the effect of energy radiation. If it is desired to keep the costs of fillers low, only a partial amount of filler is mixed in with the polymeric base material (DE-OS 2,936,926). In particular, a high-contrast light-colored inscription in the natural color of the plastic on a black background can be produced on the plastic surface by laser irradiation after incorporating 0.08 to 0.125% carbon black or graphite as filler. The whiteness of the characters can be improved
10 by optical brighteners which are added to the pigments and which are not destroyed by the laser light (German patent No. 3,044,722).

Also known are molded articles consisting of thermoplastics which contain a polymer having aromatic structure, which exhibit very good black coloring due to laser irradiation even without an additive which can change color. However, as the various grades of thermoplastics react very differently, there must be considerable coordination of process parameters, and possibly also of material modifications. It is also important to have color which is adjusted optimally for the laser light and of which the light stability remains in the usual limits for the product class concerned. In the case of large molded articles, restrictions in processing are encountered, depending on the grade of thermoplastic (Kunststoffe 78
15 (1988), issue 8, pages 688 to 691).

20 A material which can be inscribed well by laser and which has good thermal stability and stress cracking resistance is obtained by mixing polycarbonate with 10 to 50% of an aromatic polyester (European Patent No. 0,249,082).

Finally, a process for the laser inscription of high-molecular weight organic materials is known in which good inscription properties are achieved by incorporating additives (European Patent No. 0,190,997). In particular, commercially available pigments and/or polymer-soluble dyes are used as coloring additives. An inscription of adequate contrast on the surface of the materials of which the color has been adjusted with these additives can be produced by means of a
25 Nd-YAG laser preferably of doubled frequency (wavelength 532 nm after frequency doubling).

Consequently, according to the prior art, a polymeric composition can be adjusted in such a way that it can be inscribed by laser light, whether by choosing a grade of thermoplastic having good laser inscription properties or by incorporating an additive which changes color under the effect of laser irradiation.

However, in most cases the amount of color contrast which is achieved by known methods is not as high as desired.
30 Accordingly, there is a constant need for additives which can cause significant color changes to occur in the polymeric materials in which it is incorporated. Moreover, it is highly desirable that these additives not deleteriously affect the beneficial physical properties of the polymers.

SUMMARY OF THE INVENTION

35 The present invention provides a laser markable thermoplastic composition comprising at least one thermoplastic resin composition suitable for laser marking and at least one copper salt selected from copper phosphate, copper sulfate, cupric hydroxide phosphate and copper thiocyanate. The copper phosphate salt is used in an amount sufficient to cause the laser markable thermoplastic composition to absorb laser light outside the visible spectrum. Those portions of the
40 composition which are so exposed have a visibly distinct and separately identifiable color as compared to the base polymer composition.

DETAILED DESCRIPTION OF THE INVENTION

45 Most plastic resins may be effectively lasermarked through the inclusion of the copper salts according to the present invention. For example, the present invention may employ virtually any plastic resin which has a base color that allows for effective laser marking of the resin and which is not significantly degraded by the copper salts or which does not require such high processing temperatures that the copper salts are degraded during processing of the resin. According to these criteria, and depending upon the specific processing parameters, the plastic material may comprise either
50 synthetic or modified natural materials, e.g., cellulose derivatives and plastics which are obtained by polymerization, polycondensation, and polyaddition. At least the following plastic resins, including blends, copolymers and composites thereof can be suitable for lasermarking: polyolefins, polycarbonates, polyesters, rubber modified monovinylidene aromatic resins, polyetherimides, polyamides, polyester carbonates, polyphenylene sulfides, polyamideimides, polyetheramides, polyether esters, polyetherimide esters, polyarylates, polymethylpentenes, polysulfones, polyethersulfones, polystyrenes, rubber modified high impact polystyrenes, acetyls, styrene maleic anhydride copolymers, acrylonitrile styrene acrylate copolymers, polyphenylene ethers, polyether ketones, chlorinated polymers, fluorinated polymers, and liquid crystal polymers.

Thermoplastic resins are the most preferred plastic resins and, given the light base color and relatively mild processing conditions, polyesters, polycarbonates and rubber modified monovinylidene aromatic resins and their blends are particularly well suited for laser marking according to the present invention.

Through their extensive research, Applicants discovered that there are several copper salts which are surprisingly well suited for laser marking of plastic resins. In particular, the copper salts include copper II (cupric) phosphate, copper sulfate, cupric hydroxide phosphate and copper I (cuprous) thiocyanate. Applicants also discovered that a number of copper salts were not suited for laser marking of plastic resins. For example, because of their high volatility, etc., copper salts such as copper acetate, copper naphthenate and copper acetylacetonate were deemed to be unsuitable for laser marking. Similarly, other copper salts such as copper iodide, copper sulphide and copper oxide were also discovered not to be suitable for laser marking because they are black in color and/or because they do not provide adequate color contrast after exposure to laser light.

In addition to the copper salts of the invention, it may be convenient to add an additional colorant or mixture of colorants to the plastic resin. The colorant or mixture of colorants may, however, only be added in such a concentration that the laser marking produced in the practice of this invention is not impaired. Depending on the plastic resin the concentration is typically between about 0.01 to about 5% by weight.

Suitable additional colorants are inorganic or organic pigments as well as polymer-soluble dyes.

Examples of inorganic pigments are white pigments such as titanium dioxides (anatas, rutile), zinc oxide, antimony oxide, zinc sulfide, lithopones, basic lead carbonate, basic lead sulfate or basic lead silicate, and also colored pigments such as iron oxides, nickel antimony titanate, chromium antimony titanate, manganese blue, manganese violet, cobalt blue, cobalt chromium blue, cobalt nickel grey or ultramarine blue, Berlin blue, lead chromates, lead sulfochromates, molybdate orange, molybdate red, cadmium sulfides, antimony trisulfide, zirconium silicates such as zirconium vanadium blue and zirconium preseodyme yellow, and also carbon black or graphite in low concentration, and also other pigments such as aluminum pigments or mixed phase pigments in platelet form, e.g., iron oxide in platelet form doped with Al_2O_3 and/or Mn_2O_3 , as well as pearlescent pigments such as basic lead carbonate, bismuth oxychloride, bismuth oxychloride on carrier and, in particular, the titanium dioxide-coated mica pigments, which last mentioned pigments may also contain other colored metal oxides such as iron oxides, cobalt oxides, manganese oxides or chromium oxides.

Examples of organic pigments are azo, azomethine, methine, anthraquinone, indanthrone, pyranthrone, flavanthrone, benzanthrone, phthalocyanine, perinone, perylene, dioxazine, thioindigo, isoindoline, isoindolinone, quinacridone, pyrrolopyrrole or quinophthalone pigments, and also metal complexes, for example of azo, azomethine or methine dyes or metal salts of azo compounds as well as organic pigments in platelet form.

Suitable polymer-soluble dyes are, for example, disperse dyes such as those of the anthraquinone series, for example hydroxyanthraquinones, aminoanthraquinones, alkylaminoanthraquinones, hydroxyaminoanthraquinones or phenymercaptoanthraquinones, as well as metal complexes of azo dyes, in particular 1:2 chromium or cobalt complexes of monoazo dyes, and fluorescent dyes such as those of the coumarin, naphthalimide, pyrazoline, acridine, xanthene, thioxanthene, oxazine, thiazine or benzthiazole series.

In the practice of this invention, the inorganic or organic pigments or polymer-soluble dyes can be used singly or as mixtures, conveniently with or without pigment additives.

Suitable pigment additives are typically fatty acids of at least 12 carbon atoms, for example stearic acid or behenic acid and the amides, salts or esters thereof such as magnesium stearate, zinc stearate, aluminum stearate or magnesium behenate, and also quaternary ammonium compounds such as tri($\text{C}_1\text{-C}_4$)alkylbenzylammonium salts, waxes such as polyethylene wax, resin acids such as abietic acid, colophonium soap, hydrogenated or dimerised colophonium, C_{12} -18-paraffin disulfonic acids or alkylphenols, alcohols such as TCD-Alcohol M®, or vicinal aliphatic 1,2-diols.

The preparation of the plastic resins is effected by methods which are known per se, for example by incorporating the necessary colored components (molybdenum disulfide and an optional additional colorant) which may be in the form of a masterbatch, into the substrates using extruders, roll mills, mixing or grinding machines. The resultant resin is then brought into the desired final form by methods which are known per se, for example calendering, molding, extruding, coating, casting or by injection molding. It is often desirable to incorporate plasticizers into the organic material before-processing in order to produce non-brittle moldings or to diminish their brittleness. Suitable plasticizers are, for example, esters of phosphoric acid, of phthalic acid or of sebacic acid. The plasticizers may be incorporated before or after working coloring components into the polymers.

Depending on the end use, further modifiers may be added to the plastic resins, for example fillers such as kaolin, mica, feldspar, wollastonite, aluminum silicate, barium sulfate, calcium sulfate, chalk, calcite and dolomite, as well as light stabilizers, antioxidants, flame retardants, heat stabilizers, glass fibres or processing auxiliaries conventionally employed in the processing of plastics and known to the skilled person.

The thermoplastic resin composition may also include other additives which are well known in the art. For example, the resin composition may contain external lubricants, antioxidants, flame retardants or the like. If desired, ultraviolet stabilizers, flow aids, metal additives for electromagnetic radiation shielding such as nickel coated graphite fibers, anti static agents, coupling agents such as amino silanes and the like may also be added.

Energy-rich pulsed laser sources are used for marking the plastic resins according to the practice of this invention. The procedure comprises applying the radiation energy, in conformity with the shape of the marking which is to be applied, conveniently at a steep angle to the surface of the material to be marked, and focusing said radiation energy such that visibly distinct and separately identifiable marking is produced at the areas of impact without the surface of the marked material being perceptibly damaged.

Examples of such energy source are solid state pulsed lasers such as ruby lasers or frequency multiplied Nd:YAG lasers, pulsed lasers with booster such as pulsed dye lasers or Raman shifter, and also continuous wave lasers with pulse modifications (Q-switch, mode locker), for example on the basis of CW Nd:YAG lasers with frequency multiplier, or CW ion lasers (Ar, Kr), as well as pulsed metal vapor lasers, for example copper vapor lasers or gold vapor lasers, or high capacity pulsed semi-conductor lasers which emit visible light by frequency doubling, and also pulsed gas lasers such as excimer and nitrogen lasers.

Depending on the laser system employed, pulse contents of up to several Joules per cm², intensities of up to 10¹² W/cm², pulse durations of from 10⁻¹⁵ seconds to 10⁻⁶ seconds and frequencies of up to 10⁹ Hz are possible. Pulse contents of micro-Joule to kilo-Joule, intensities of kilowatt/cm² to 100 megawatt/cm², pulse durations of microseconds to picoseconds, and frequencies of a few hertz to 50 kilohertz are advantageously used.

Preferred lasers are pulsed or pulse-modified, frequency doubled Nd:YAG lasers or metal vapor lasers such as gold or, in particular, copper vapor lasers, as well as excimer lasers.

The following table lists a number of commercially available lasers which may be suitably used in the practice of this invention.

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TABLE

Type/Representative	Examples of commercially available types	Principal wavelength (subsidiary wavelengths) [nm]
Solid state pulsed lasers		
Ruby laser	Lasermetrics (938R6R4L-4)	694 (347)
Nd:YAG laser	Quanta Ray (DCR 2A)	1064, (532, 355, 266)
Alexandrite laser	Apollo (7562)	730-780
Pulsed lasers with booster such as		
Raman shifter	Quanta Ray (RS-1)	UV-IR
Dye laser	Lambda Physik FL 2002	ca. 300-1000
CW laser with pulse modification		
ND:YAG	Lasermetrics	532
(Q-Switch, 2ω)	(9560QTG)	
Argon (mode-locked)	Spectra-Physics SP 2030	514.5, 488
Pulsed metal vapor laser		
Cu vapor laser	Plasma-Kinetics 751	510, 578
Au vapor laser	Plasma-Kinetics	628
Mn vapor laser	Oxford	534, 1290,
Pb vapor laser	Laser CU 25	723
Semi-conductor diode lasers	M/ACOM Type LD 65	ca. 905, (402)
Semi-conductor diode laser array	STANTEL Type LF 100	ca. 905 (402)
Pulsed gas lasers Excimer		
XeCl	Lambda Physik	308
XeF as well as	EMG-103	351
N ₂		337

Lasers whose parameters can be readily adjusted, for example pulse content and pulse duration, permit the best possible adaptation to the requirements of the materials to be marked.

The best wavelength to be selected for the irradiation is that at which the radiation-sensitive copper salts and the optional additional colorant absorbs most strongly, and that at which the plastics material to be marked absorbs little.

Preferably laser light with a wavelength in the IR range is used. The "near IR range" as meaning the range from about 0.78 μ m to about 2 μ m.

Three different methods are normally suitable for laser marking in the practice of this invention: the mask method, the linear marking method and the dot matrix method. In these last two mentioned methods (dynamic focusing), the laser is preferably combined with a laser marking system, so that the plastics material can be marked with any, e.g. computer-programmed, digits, letters and special symbols.

The choice of laser system in respect to capacity and frequency depends basically on the marking method employed. The high capacity and low frequency of, e.g., solid state pulsed lasers and excimer lasers are preferred for mask exposure. The average to low capacities and rapid frequencies of pulsed metal vapor lasers or of continuous wave lasers with pulse modifications are preferred for producing markings that require dynamic focusing. Beam deflection can be effected, for example, acousto-optically, holographically, with galvo-mirrors or polygon scanners. Dynamic focusing makes possible an extremely flexible marking, as the marks can be produced electronically.

EXAMPLE 1

This example demonstrates several of the copper salts which were found to provide effective laser marking of thermoplastic resins. The copper compounds were evaluated in a thermoplastic polyester sold by General Electric Co. as Valox® 325C, a grade specially developed for keycap applications.

A Nd/YAG laser, operating at a wavelength of 1064nm was used to mark the disks. Apart from the material composition, the laser settings were of relatively major importance. The optimal settings were dependent on scan speed, lamp current, Q-switch and mode blender. To determine the optimal settings, the lamp current and Q-switch were varied as function of the scan speed. Former results indicated that the mode blender should have a diaphragm of 1.0 mm. The settings providing the best contrast were used to mark a small square (1*1cm). This area allowed a measurement of color differences of the background and the marked area (CieLab method, DIN 6174, source D65). A Delta E value of 20 is generally considered as sufficient.

30

A. Copper Sulfate						
	1	2	3	4	5	6
Valox® 325C	100	100	100	100	100	100
Titanium dioxide	2.8	4.3	4.3	4.3	4.3	4.3
Carbon black	0.015	0.015	0.015	0.015	0.015	0.015
Copper sulfate (solid)		1.5				
Copper sulfate (solution)			0.05	0.1	0.5	1.5
Delta E	12.5	16.7	21.3	21.4	19.1	14.8

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B. Copper Phosphate						
	1	7	8	9	10	11
Valox® 325C	100	100	100	100	100	100
Titanium dioxide	2.8	4.3	4.3	4.3	4.3	4.3
Carbon black	0.015	0.015	0.015	0.015	0.015	0.015
Copper phosphate (PS=30 µm)		3				
Copper phosphate (PS=15 µm)			3	1.5		
Copper phosphate (PS=6 µm)					3	1.5
Delta E	12.5	17.7	23.6	20.6	25.4	24.1

⁵ 4PS=particle size, 90% of particles is smaller than given number.

C. Copper Thiocyanate					
	1	12	13	14	15
Valox® 325C	100	100	100	100	100
Titanium dioxide	2.8	4.3	4.3	4.3	4.3
Carbon black	0.015	0.015	0.015	0.015	0.015
Copper thiocyanate (PS=0.4 µm)		0.75	1.0	1.5	3
Delta E	12.5	20	21.9	23.3	19.9

D. Copper Thiocyanate			
	16	17	18
Valox® 325C	100	100	100
Titanium dioxide	2.8	4.3	4.3
Copper thiocyanate (PS=0.4 µm)		1.5	3
Delta E	25.5	36.6	38.5

⁵ Addition of 3 pbw of copper thiocyanate to a Valox® 325C formulation, improved the sensibility for the CO₂-laser significantly. Without the laser sensitive additive, the plastic material was just ablated, while in the case where the additive was present, a grey-brown symbol appeared after irradiation.

EXAMPLE 2

⁵⁵ A direct comparison was carried out between Cupric phosphate, Cuprous thiocyanate and Cupric hydroxide phosphate as possible contrast enhancing additives.

The following additives were evaluated in a polyester resin manufactured by General Electric and sold under the trademark Valox® 325C.

Additives used:

Cupric hydroxide phosphate; Aldrich 34,440-0, 97%.

Cupric phosphate; Schmidt B.V. M13-3²

Cuprous thiocyanate; HCA Holland Colors

5 All laser experiments were carried out using a Nd:YAG 1064 nm laser. Model LBI 6000. Supplier; Carl Baasel lasertechnik GmbH. Focus: 160 mm, Working distance: 180 mm.

Laser settings were optimized in the same way as Example 1. Contrast has been measured and defined as delta E (marked area versus background) according to CieLab method, DIN 6174, source D65.

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A. Cupric hydroxide phosphate						
Cu ₂ (OH)PO ₄ (pbw)	0	0.5	1	1.5	2	3
(composition in pbw)						
Valox® 325C	100	100	100	100	100	100
Titanium dioxide	4.3	4.3	4.3	4.3	4.3	4.3
Carbon Black	0.015	0.015	0.015	0.015	0.015	0.015
(laser settings)						
Modeblendor (mm)	1.0	1.0	1.0	1.0	1.0	1.0
Lampcurrent (Ampere)	18	16	14	14	14	14
Q-switch (Herz)	2000	2000	2000	2000	2000	2000
Speed (mm/s)	200	175	225	225	225	225
Wobble (mm)	0	0	0	0	0	0
delta E	16.3	15.8	18.9	20.4	23.2	21.7

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B. Cupric phosphate						
Cu ₃ (PO ₄) ₂ (pbw)	0	0.5	1	1.5	2	3
(composition in pbw)						
Valox® 325C	100	100	100	100	100	100
Titanium dioxide	4.3	4.3	4.3	4.3	4.3	4.3
Carbon Black	0.015	0.015	0.015	0.015	0.015	0.015
(laser settings)						
Modeblendor (mm)	1.0	1.0	1.0	1.0	1.0	1.0
Lampcurrent (Ampere)	16	16	14	14	14	14
Q-switch (Herz)	2000	2000	2000	2000	3000	3000
Speed (mm/s)	225	200	200	200	225	225
Wobble (mm)	0	0	0	0	0	0
delta E	16.3	17.2	18.8	20.8	20.7	22.6

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C. Cuprous thiocyanate					
CuSCN (pbw)	0	0.5	1	1.5	
(composition in pbw)					
Valox® 325C	100	100	100	100	
Titanium dioxide	4.3	4.3	4.3	4.3	
Carbon Black	0.015	0.015	0.015	0.01	
(lasersettings)					
Modeblendor (mm)	1.0	1.0	1.0	1.0	
Lampcurrent (Ampere)	16	16	14	14	
Q-switch (Herz)	2000	2000	2000	2000	
Speed (mn/s)	225	225	225	225	
Wobble (mm)	0	0	0	0	
delta E	16.3	18.8	22.3	24.7	

Obviously, other modifications and variations of the present invention are possible in light of the above teachings. It is, therefore, to be understood that changes may be made in the particular embodiments described above which are within the scope of the invention as defined in the appended claims.

Claims

1. A laser markable thermoplastic composition, comprising:
 - (a) at least one thermoplastic resin composition suitable for laser marking; and
 - (b) at least one copper salt selected from the group consisting of copper phosphate, copper sulfate, cupric hydroxide phosphate and copper thiocyanate, wherein said copper salt is present in an amount sufficient to cause said laser markable thermoplastic composition to absorb a laser light outside the visible spectrum such that the portion of said composition which absorbs said laser light has a visibly distinct and separately identifiable color.
2. The thermoplastic composition of Claim 1, wherein the wavelength of said laser light is above about 900nm.
3. The thermoplastic composition of Claim 1, wherein said copper salt is copper phosphate or copper thiocyanate.
4. The thermoplastic composition of Claim 1 wherein said thermoplastic resin comprises at least one resin selected from polycarbonate, polyester, rubber modified monovinyldene aromatics, polyetherimide, polyesteramides, polyamide, polyester carbonates, polyphenylene sulfide, polyamideimide, polyether esters, polyetherimide esters, polycarbonate, polymethylpentene, polysulfone polyethersulfone, polystyrene, rubber modified high impact polystyrene acetyl, polyphenylene ether, polyether ketone, chlorinated polymer, fluorinated polymer, liquid crystal polymer, copolymers of the above, or blends of the above.
5. The thermoplastic composition of Claim 4, wherein said thermoplastic resin comprises polyester, polycarbonate, polyestercarbonate, rubber modified monovinyldene aromatics, polyetherimides, polyesteramides, polyetherester, polyetherimide ester, or copolymers or blends of the above.
6. The thermoplastic composition of Claim 5, wherein said thermoplastic resins comprises polycarbonate polyalkylene terephthalate, rubber modified monovinyldene aromatics, or copolymers or blends of the above.
7. The thermoplastic composition of Claim 2, wherein the amount of said copper salt ranges from about 0.05 to about 10 parts by weight of the thermoplastic composition.

8. The thermoplastic composition of Claim 7, wherein the amount of said copper salt ranges from about 0.1 to about 5 parts by weight of the thermoplastic composition.
- 5 9. The thermoplastic composition of Claim 1, wherein the amount of said copper salt ranges from about 0.5 to about 3 parts by weight of the thermoplastic composition.
- 10 10. The thermoplastic composition of Claim 2, wherein the particle size of said copper salt is less than about 10 µm.
11. The thermoplastic composition of Claim 10, wherein the particle size of said copper salt is less than about 1 µm.
- 10 12. The thermoplastic composition of Claim 2, wherein the total color difference between the base composition and the portion which absorbed said laser light is at least 10 units as measured by the Delta E.
- 15 13. The thermoplastic composition of Claim 12, wherein the total color difference between the base composition and the portion which absorbed said laser light is at least 15 units as measured by the Delta E.
14. The thermoplastic composition of Claim 13, wherein the total color difference between the base composition and the portion which absorbed said laser light is at least 20 units as measured by the Delta E.
- 20 15. The thermoplastic composition of Claim 2, wherein said composition further comprises carbon black or titanium dioxide.
16. A laser markable article of manufacture comprising the composition of Claim 1.
- 25 17. A process for producing a laser markable thermoplastic composition comprising extruding the composition of Claim 1.
18. A process for producing a laser marked article of manufacture comprising the steps of:
 - 30 (I) extruding and molding a thermoplastic composition comprising:
 - (a) at least one thermoplastic resin composition suitable for laser marking; and
 - (b) at least one copper salt selected from the group consisting of copper phosphate, copper sulfate, cupric hydroxide phosphate or copper thiocyanate, wherein said copper salt is present in an amount sufficient to cause said laser markable thermoplastic composition to absorb laser light outside the visible spectrum such that the portion of said composition which absorbs said laser light has a visibly distinct and separately identifiable color; and
 - 35 (II) laser marking said article with a laser having a wavelength outside the visible spectrum.

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European Patent
Office

EUROPEAN SEARCH REPORT

Application Number
EP 95 10 3734

DOCUMENTS CONSIDERED TO BE RELEVANT			
Category	Citation of document with indication, where appropriate, of relevant passages	Relevant to claim	CLASSIFICATION OF THE APPLICATION (Int.Cl.6)
E	EP-A-0 665 118 (HUELS CHEMISCHE WERKE AG) 2 August 1995 * the whole document * ---	1-9	C08K3/30 C08K3/32 B41M5/24
X	EP-A-0 542 115 (BASF AG) 19 May 1993 * page 8, line 50 - page 9, line 38 * * claims 1-4 * ---	1,2,4-9, 15,16	
X	EP-A-0 400 305 (HUELS CHEMISCHE WERKE AG) 5 December 1990 * examples * * claims 1-6 * -----	1,2,4-9, 15,16	
TECHNICAL FIELDS SEARCHED (Int.Cl.6)			
C08K			
The present search report has been drawn up for all claims			
Place of search	Date of completion of the search	Examiner	
THE HAGUE	29 November 1995	Siemens, T	
CATEGORY OF CITED DOCUMENTS		T : theory or principle underlying the invention E : earlier patent document, but published on, or after the filing date D : document cited in the application L : document cited for other reasons ----- & : member of the same patent family, corresponding document	
X : particularly relevant if taken alone Y : particularly relevant if combined with another document of the same category A : technological background O : non-written disclosure P : intermediate document			



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Kupfersalze zur Lasermarkierung von thermoplastischen Zusammensetzungen

Sels de cuivre pour marquage par laser de compositions thermoplastiques

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(56) References cited:
EP-A- 0 400 305 EP-A- 0 542 115
EP-A- 0 665 118

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Description**BACKGROUND OF THE INVENTION**

- 5 [0001] It is known to make a polymeric material which can be inscribed by means of laser light by mixing in a filler which changes color under the effect of energy radiation. If it is desired to keep the costs of fillers low, only a partial amount of filler is mixed in with the polymeric base material (DE-OS 2,936,926). In particular, a high-contrast light-colored inscription in the natural color of the plastic on a black background can be produced on the plastic surface by laser irradiation after incorporating 0.08 to 0.125% carbon black or graphite as filler. The whiteness of the characters
10 can be improved by optical brighteners which are added to the pigments and which are not destroyed by the laser light (German patent No. 3,044,722).
- [0002] Also known are molded articles consisting of thermoplastics which contain a polymer having aromatic structure, which exhibit very good black coloring due to laser irradiation even without an additive which can change color. However, as the various grades of thermoplastics react very differently, there must be considerable coordination of
15 process parameters, and possibly also of material modifications. It is also important to have color which is adjusted optimally for the laser light and of which the light stability remains in the usual limits for the product class concerned. In the case of large molded articles, restrictions in processing are encountered, depending on the grade of thermoplastic (Kunststoffe 78 (1988), issue 8, pages 688 to 691).
- 20 [0003] A material which can be inscribed well by laser and which has good thermal stability and stress cracking resistance is obtained by mixing polycarbonate with 10 to 50% of an aromatic polyester (European Patent No. 0,249,082).
- [0004] Finally, a process for the laser inscription of high-molecular weight organic materials is known in which good inscription properties are achieved by incorporating additives (European Patent No. 0,190,997). In particular, commercially available pigments and/or polymer-soluble dyes are used as coloring additives. An inscription of adequate contrast on the surface of the materials of which the color has been adjusted with these additives can be produced by means of a Nd-YAG laser preferably of doubled frequency (wavelength 532 nm after frequency doubling).
- 25 [0005] Consequently, according to the prior art, a polymeric composition can be adjusted in such a way that it can be inscribed by laser light, whether by choosing a grade of thermoplastic having good laser inscription properties or by incorporating an additive which changes color under the effect of laser irradiation.
- 30 [0006] However, in most cases the amount of color contrast which is achieved by known methods is not as high as desired. Accordingly, there is a constant need for additives which can cause significant color changes to occur in the polymeric materials in which it is incorporated. Moreover, it is highly desirable that these additives not deleteriously affect the beneficial physical properties of the polymers.

SUMMARY OF THE INVENTION

- 35 [0007] The present invention provides a laser markable thermoplastic composition comprising at least one thermoplastic resin composition suitable for laser marking and at least one copper salt selected from copper phosphate, copper sulfate, and copper thiocyanate wherein said copper salt is present in an amount of 0.05 to 10 parts by weight per 100 parts by weight of the thermoplastic resin composition. The copper phosphate salt is used in an amount sufficient to cause the laser markable thermoplastic composition to absorb laser light outside the visible spectrum. Those portions of the composition which are so exposed have a visibly distinct and separately identifiable color as compared to the base polymer composition.

DETAILED DESCRIPTION OF THE INVENTION

- 40 [0008] Most plastic resins may be effectively lasermarked through the inclusion of the copper salts according to the present invention. For example, the present invention may employ virtually any plastic resin which has a base color that allows for effective laser marking of the resin and which is not significantly degraded by the copper salts or which does not require such high processing temperatures that the copper salts are degraded during processing of the resin.
45 According to these criteria, and depending upon the specific processing parameters, the plastic material may comprise either synthetic or modified natural materials, e.g., cellulose derivatives and plastics which are obtained by polymerization, polycondensation, and polyaddition. At least the following plastic resins, including blends, copolymers and composites thereof can be suitable for lasermarking: polyolefins, polycarbonates, polyesters, rubber modified monovinylidene aromatic resins, polyetherimides, polyamides, polyester carbonates, polyphenylene sulfides, polyamideimides, polyesteramides, polyether esters, polyetherimide esters, polyarylates, polymethylpentenes, polysulfones, polyethersulfones, polystyrenes, rubber modified high impact polystyrenes, styrene maleic anhydride copolymers, acrylonitrile styrene acrylate copolymers, polyphenylene ethers, polyether ketones, chlorinated polymers, fluorinated poly-

mers, and liquid crystal polymers.

[0009] Thermoplastic resins are the most preferred plastic resins and, given the light base color and relatively mild processing conditions, polyesters, polycarbonates and rubber modified monovinylidene aromatic resins and their blends are particularly well suited for laser marking according to the present invention.

[0010] Through their extensive research, Applicants discovered that there are several copper salts which are surprisingly well suited for laser marking of plastic resins. In particular, the copper salts include copper II (cupric) phosphate, copper sulfate and copper I (cuprous) thiocyanate. Applicants also discovered that a number of copper salts were not suited for laser marking of plastic resins. For example, because of their high volatility, etc., copper salts such as copper acetate, copper naphthenate and copper acetylacetone were deemed to be unsuitable for laser marking. Similarly, other copper salts such as copper iodide, copper sulphide and copper oxide were also discovered not to be suitable for laser marking because they are black in color and/or because they do not provide adequate color contrast after exposure to laser light.

[0011] In addition to the copper salts of the invention, it may be convenient to add an additional colorant or mixture of colorants to the plastic resin. The colorant or mixture of colorants may, however, only be added in such a concentration that the laser marking produced in the practice of this invention is not impaired. Depending on the plastic resin the concentration is typically between about 0.01 to about 5% by weight.

[0012] Suitable additional colorants are inorganic or organic pigments as well as polymer-soluble dyes.

[0013] Examples of inorganic pigments are white pigments such as titanium dioxides (anatas, rutile), zinc oxide, antimony oxide, zinc sulfide, lithopones, basic lead carbonate, basic lead sulfate or basic lead silicate, and also colored pigments such as iron oxides, nickel antimony titanate, chromium antimony titanate, manganese blue, manganese violet, cobalt blue, cobalt chromium blue, cobalt nickel grey or ultramarine blue, Berlin blue, lead chromates, lead sulfochromates, molybdate orange, molybdate red, cadmium sulfides, antimony trisulfide, zirconium silicates such as zirconium vanadium blue and zirconium praseodyme yellow, and also carbon black or graphite in low concentration, and also other pigments such as aluminum pigments or mixed phase pigments in platelet form, e.g., iron oxide in platelet form doped with Al_2O_3 and/or Mn_2O_3 , as well as pearlescent pigments such as basic lead carbonate, bismuth oxychloride, bismuth oxychloride on carrier and, in particular, the titanium dioxide-coated mica pigments, which last mentioned pigments may also contain other colored metal oxides such as iron oxides, cobalt oxides, manganese oxides or chromium oxides.

[0014] Examples of organic pigments are azo, azomethine, methine, anthraquinone, indanthrone, pyranthrone, flavanthrone, benzanthrone, phthalocyanine, perinone, perylene, dioxazine, thioindigo, isoindoline, isoindolinone, quinacridone, pyrrolopyrrole or quinophthalone pigments, and also metal complexes, for example of azo, azomethine or methine dyes or metal salts of azo compounds as well as organic pigments in platelet form.

[0015] Suitable polymer-soluble dyes are, for example, disperse dyes such as those of the anthraquinone series, for example hydroxyanthraquinones, aminoanthraquinones, alkylaminoanthraquinones, hydroxyaminoanthraquinones or phenylmercaptoanthraquinones, as well as metal complexes of azo dyes, in particular 1:2 chromium or cobalt complexes of monoazo dyes, and fluorescent dyes such as those of the coumarin, naphthalimide, pyrazoline, acridine, xanthene, thioxanthene, oxazine, thiazine or benzthiazole series.

[0016] In the practice of this invention, the inorganic or organic pigments or polymer-soluble dyes can be used singly or as mixtures, conveniently with or without pigment additives.

[0017] Suitable pigment additives are typically fatty acids of at least 12 carbon atoms, for example stearic acid or behenic acid and the amides, salts or esters thereof such as magnesium stearate, zinc stearate, aluminum stearate or magnesium behenate, and also quaternary ammonium compounds such as tri(C₁-C₄)alkylbenzylammonium salts, waxes such as polyethylene wax, resin acids such as abietic acid, colophonum soap, hydrogenated or dimerised colophonum, C₁₂-C₁₈-paraffin disulfonic acids or alkylphenols, alcohols such as TCD-Alcohol M®, or vicinal aliphatic 1,2-diols.

[0018] The preparation of the plastic resins is effected by methods which are known per se, for example by incorporating the necessary colored components (molybdenum disulfide and an optional additional colorant) which may be in the form of a masterbatch, into the substrates using extruders, roll mills, mixing or grinding machines. The resultant resin is then brought into the desired final form by methods which are known per se, for example calendering, molding, extruding, coating, casting or by injection molding. It is often desirable to incorporate plasticizers into the organic material before-processing in order to produce non-brittle moldings or to diminish their brittleness. Suitable plasticizers are, for example, esters of phosphoric acid, of phthalic acid or of sebacic acid. The plasticizers may be incorporated before or after working coloring components into the polymers.

[0019] Depending on the end use, further modifiers may be added to the plastic resins, for example fillers such as kaolin, mica, feldspar, wollastonite, aluminum silicate, barium sulfate, calcium sulfate, chalk, calcite and dolomite, as well as light stabilizers, antioxidants, flame retardants, heat stabilizers, glass fibres or processing auxiliaries conventionally employed in the processing of plastics and known to the skilled person.

[0020] The thermoplastic resin composition may also include other additives which are well known in the art. For

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example, the resin composition may contain external lubricants, antioxidants, flame retardants or the like. If desired, ultraviolet stabilizers, flow aids, metal additives for electromagnetic radiation shielding such as nickel coated graphite fibers, anti static agents, coupling agents such as amino silanes and the like may also be added.

[0021] Energy-rich pulsed laser sources are used for marking the plastic resins according to the practice of this invention. The procedure comprises applying the radiation energy, in conformity with the shape of the marking which is to be applied, conveniently at a steep angle to the surface of the material to be marked, and focusing said radiation energy such that visibly distinct and separately identifiable marking is produced at the areas of impact without the surface of the marked material being perceptibly damaged.

[0022] Examples of such energy source are solid state pulsed lasers such as ruby lasers or frequency multiplied Nd:YAG lasers, pulsed lasers with booster such as pulsed dye lasers or Raman shifter, and also continuous wave lasers with pulse modifications (Q-switch, mode locker), for example on the basis of CW Nd:YAG lasers with frequency multiplier, or CW ion lasers (Ar, Kr), as well as pulsed metal vapor lasers, for example copper vapor lasers or gold vapor lasers, or high capacity pulsed semi-conductor lasers which emit visible light by frequency doubling, and also pulsed gas lasers such as excimer and nitrogen lasers.

[0023] Depending on the laser system employed, pulse contents of up to several Joules per cm², intensities of up to 10¹² W/cm², pulse durations of from 10⁻¹⁵ seconds to 10⁻⁶ seconds and frequencies of up to 10⁹ Hz are possible. Pulse contents of micro-Joule to kilo-Joule, Intensities of kilowatt/cm² to 100 megawatt/cm², pulse durations of microseconds to picoseconds, and frequencies of a few hertz to 50 kilohertz are advantageously used.

[0024] Preferred lasers are pulsed or pulse-modified, frequency doubled Nd:YAG lasers or metal vapor lasers such as gold or, in particular, copper vapor lasers, as well as excimer lasers.

[0025] The following table lists a number of commercially available lasers which may be suitably used in the practice of this invention.

TABLE

Type/Representative	Examples of commercially available types	Principal wavelength (subsidiary wavelengths) [nm]
Solid state pulsed lasers		
Ruby laser	Lasermetrics (938R6R4L-4)	694 (347)
Nd:YAG laser	Quanta Ray (DCR 2A)	1064, (532, 355, 266)
Alexandrite laser	Apollo (7562)	730-780
Pulsed lasers with booster such as		
Raman shifter	Quanta Ray (RS-1)	UV-IR
Dye laser	Lambda Physik FL 2002	ca. 300-1000
CW laser with pulse modification		
ND:YAG (Q-Switch, 2ω)	Lasermetrics (9560QTG)	532
Argon (mode-locked)	Spectra-Physics SP 2030	514.5, 488
Pulsed metal vapor laser		
Cu vapor laser	Plasma-Kinetics 751	510, 578
Au vapor laser	Plasma-Kinetics	628
Mn vapor laser	Oxford	534, 1290,
Pb vapor laser	Laser CU 25	723
Semi-conductor diode lasers	M/ACOM Type LD 65	ca. 905, (402)
Semi-conductor diode laser array	STANTEL Type LF 100	ca. 905 (402)
Pulsed gas lasers Excimer		
XeCl	Lambda Physik	308
XeF as well as	EMG-103	351

TABLE (continued)

Type/Representative	Examples of commercially available types	Principal wavelength (subsidiary wavelengths) [nm]
Pulsed gas lasers Excimer		
N ₂		337

[0026] Lasers whose parameters can be readily adjusted, for example pulse content and pulse duration, permit the best possible adaptation to the requirements of the materials to be marked.

[0027] The best wavelength to be selected for the irradiation is that at which the radiation-sensitive copper salts and the optional additional colorant absorbs most strongly, and that at which the plastics material to be marked absorbs little.

[0028] Preferably laser light with a wavelength in the IR range is used. The "near IR range" as meaning the range from about 0.78μm to about 2μm.

[0029] Three different methods are normally suitable for laser marking in the practice of this Invention: the mask method, the linear marking method and the dot matrix method. In these last two mentioned methods (dynamic focusing), the laser is preferably combined with a laser marking system, so that the plastics material can be marked with any, e.g. computer-programmed, digits, letters and special symbols.

[0030] The choice of laser system in respect to capacity and frequency depends basically on the marking method employed. The high capacity and low frequency of, e.g., solid state pulsed lasers and excimer lasers are preferred for mask exposure. The average to low capacities and rapid frequencies of pulsed metal vapor lasers or of continuous wave lasers with pulse modifications are preferred for producing markings that require dynamic focusing. Beam deflection can be effected, for example, acousto-optically, holographically, with galvo-mirrors or polygon scanners. Dynamic focusing makes possible an extremely flexible marking, as the marks can be produced electronically.

EXAMPLE 1

[0031] This example demonstrates several of the copper salts which were found to provide effective laser marking of thermoplastic resins. The copper compounds were evaluated in a thermoplastic polyester sold by General Electric Co. as Valox® 325C, a grade specially developed for keycap applications.

[0032] A Nd/YAG laser, operating at a wavelength of 1064nm was used to mark the disks. Apart from the material composition, the laser settings were of relatively major importance. The optimal settings were dependent on scan speed, lamp current, Q-switch and mode blinder. To determine the optimal settings, the lamp current and Q-switch were varied as function of the scan speed. Former results indicated that the mode blinder should have a diaphragm of 1.0 mm. The settings providing the best contrast were used to mark a small square (1*1cm). This area allowed a measurement of color differences of the background and the marked area (CieLab method, DIN 6174, source D65). A Delta E value of 20 is generally considered as sufficient.

[0033] A Delta E value of at least 10 units, more preferably at least 15 units, most preferably at least 20 units as measured by the Delta E is preferred.

A. Copper Sulfate

	1	2	3	4	5	6
Valox® 325C	100	100	100	100	100	100
Titanium dioxide	2.8	4.3	4.3	4.3	4.3	4.3
Carbon black	0.015	0.015	0.015	0.015	0.015	0.015
Copper sulfate (solid)		1.5				
Copper sulfate (solution)			0.05	0.1	0.5	1.5
Delta E	12.5	16.7	21.3	21.4	19.1	14.8

B. Copper Phosphate

	1	7	8	9	10	11
Valox® 325C	100	100	100	100	100	100
Titanium dioxide	2.8	4.3	4.3	4.3	4.3	4.3
Carbon black	0.015	0.015	0.015	0.015	0.015	0.015
Copper phosphate (PS=30 µm)		3				
Copper phosphate (PS=15 µm)			3	1.5		
Copper phosphate (PS=6 µm)					3	1.5
Delta E	12.5	17.7	23.6	20.6	25.4	24.1

*PS=particle size, 90% of particles is smaller than given number.

C. Copper Thiocyanate

	1	12	13	14	15
Valox® 325C	100	100	100	100	100
Titanium dioxide	2.8	4.3	4.3	4.3	4.3
Carbon black	0.015	0.015	0.015	0.015	0.015
Copper thiocyanate (PS=0.4 µm)		0.75	1.0	1.5	3
Delta E	12.5	20	21.9	23.3	19.9

D. Copper Thiocyanate

	16	17	18
Valox® 325C	100	100	100
Titanium dioxide	2.8	4.3	4.3
Copper thiocyanate (PS ⁴ =0.4 µm)		1.5	3
Delta E	25.5	36.6	38.5

Addition of 3 pbw of copper thiocyanate to a Valox® 325C formulation, improved the sensibility for the CO₂-laser significantly. Without the laser sensitive additive, the plastic material was just ablated, while in the case where the additive was present, a grey-brown symbol appeared after irradiation.

EXAMPLE 2

[0034] A direct comparison was carried out between Cupric phosphate, Cuprous thiocyanate and Cupric hydroxide phosphate as possible contrast enhancing additives.

5 [0035] The following additives were evaluated in a polyester resin manufactured by General Electric and sold under the trademark Valox® 325C.

[0036] Additives used:

Cupric hydroxide phosphate; Aldrich 34,440-0, 97%.

10 Cupric phosphate; Scmidt B.V. M13-3²

Cuprous thiocyanate; HCA Holland Colors

[0037] All laser experiments were carried out using a Nd:YAG 1064 nm laser. Model LBI 6000. Supplier; Carl Baasel lasertechnik GmbH. Focus: 160 mm, Working distance: 180 mm.

15 [0038] Laser settings were optimized in the same way as Example 1. Contrast has been measured and defined as delta E (marked area versus background) according to CieLab method, DIN 6174, source D65.

A. Cupric hydroxide phosphate

$\text{Cu}_2(\text{OH})\text{PO}_4$ (pbw)	0	0.5	1	1.5	2	3
(composition in pbw)						
Valox® 325C	100	100	100	100	100	100
Titanium dioxide	4.3	4.3	4.3	4.3	4.3	4.3
Carbon Black	0.015	0.015	0.015	0.015	0.015	0.015
(laser settings)						
Modeblendor (mm)	1.0	1.0	1.0	1.0	1.0	1.0
Lampcurrent (Ampere)	18	16	14	14	14	14
Q-switch (Herz)	2000	2000	2000	2000	2000	2000
Speed (mm/s)	200	175	225	225	225	225
Wobble (mm)	0	0	0	0	0	0
delta E	16.3	15.8	18.9	20.4	23.2	21.7

B. Cupric phosphate

$\text{Cu}_3(\text{PO}_4)_2$ (pbw)	0	0.5	1	1.5	2	3
(composition in pbw)						
Valox® 325C	100	100	100	100	100	100
Titanium dioxide	4.3	4.3	4.3	4.3	4.3	4.3
Carbon Black	0.015	0.015	0.015	0.015	0.015	0.015
(laser settings)						
Modeblendor (mm)	1.0	1.0	1.0	1.0	1.0	1.0
Lampcurrent (Ampere)	16	16	14	14	14	14
Q-switch (Herz)	2000	2000	2000	2000	3000	3000
Speed (mm/s)	225	200	200	200	225	225
Wobble (mm)	0	0	0	0	0	0
delta E	16.3	17.2	18.8	20.8	20.7	22.6

C. Cuprous thiocyanate

CuSCN (pbw)	0	0.5	1	1.5
(composition in pbw)				
Valox® 325C	100	100	100	100
Titanium dioxide	4.3	4.3	4.3	4.3
Carbon Black	0.015	0.015	0.015	0.01
(lasersettings)				
Modeblend or (mm)	1.0	1.0	1.0	1.0
Lampcurrent (Ampere)	16	16	14	14
Q-switch (Herz)	2000	2000	2000	2000
Speed (mn/s)	225	225	225	225
Wobble (mm)	0	0	0	0
delta E	16.3	18.8	22.3	24.7

Claims

1. A laser markable thermoplastic composition, comprising:
- (a) at least one thermoplastic resin composition suitable for laser marking; and
 - (b) at least one copper salt selected from the group consisting of copper phosphate, copper sulfate, and copper thiocyanate, wherein said copper salt is present in an amount of 0.05 to 10 parts by weight per 100 parts by weight of the thermoplastic composition.
2. The thermoplastic composition of Claim 1, wherein said copper salt is copper phosphate or copper thiocyanate.
3. The thermoplastic composition of Claim 1 wherein said thermoplastic resin comprises at least one resin selected from polycarbonate, polyester, rubber modified monovinylidene aromatic resins, polyetherimide, polyesteramides, polyamide, polyester carbonates, polyphenylene sulfide, polyamideimide, polyether esters, polyetherimide esters, polyarylate, polymethylpentene, polysulfone polyethersulfone, polystyrene, rubber modified high impact polystyrene, polyphenylene ether, polyether ketone, chlorinated polymer, fluorinated polymer, liquid crystal polymer, copolymers of the above, or blends of the above.
4. The thermoplastic composition of claim 4, wherein said thermoplastic resin comprises polyester, polycarbonate, polyestercarbonate, rubber modified monovinylidene aromatics, polyetherimides, polyesteramides, polyetherester, polyetherimide ester, or copolymers or blends of the above.
5. The thermoplastic composition of Claim 5, wherein said thermoplastic resins comprises polycarbonate polyalkylene terephthalate, rubber modified monovinylidene aromatics, or copolymers or blends of the above.
6. The thermoplastic composition of Claim 6 wherein the amount of said copper salt ranges from 0.1 to 5 parts by weight of the thermoplastic composition.
7. The thermoplastic composition of Claim 1, wherein the amount of said copper salt ranges from 0.5 to 3 parts by weight of the thermoplastic composition.
8. The thermoplastic composition of Claim 2, wherein the particle size of said copper salt is less than 10 µm.
9. The thermoplastic composition of Claim 9, wherein the particle size of said copper salt is less than 1 µm.

10. The thermoplastic composition of Claim 12, wherein the total color difference between the base composition and
the portion which absorbed said laser light is at least 20 units as measured by the Delta E.

5 11. The thermoplastic composition of Claim 2, wherein said composition further comprises carbon black or titanium
dioxide.

12. A laser markable article of manufacture comprising the composition of Claim 1.

10 13. A process for producing a laser markable thermoplastic composition comprising extruding the composition of Claim
1.

14. A process for producing a laser marked article of manufacture comprising the steps of:

15 (I) extruding and molding a thermoplastic composition comprising:

(a) at least one thermoplastic resin composition suitable for laser marking; and
(b) at least one copper salt selected from the group consisting of copper phosphate, copper sulfate, or
copper thiocyanate, wherein said copper salt is present in an amount sufficient to cause said laser mark-
able thermoplastic composition to absorb laser light outside the visible spectrum such that the portion of
said composition which absorbs said laser light has a visibly distinct and separately identifiable color; and

20 (II) laser marking said article with a laser having a wavelength outside the visible spectrum.

25 15. A process according to claim 14, wherein the wave length of the laser is above 900 nm.

Patentansprüche

30 1. Eine lasermarkierbare thermoplastische Zusammensetzung aufweisend:

(a) wenigstens eine zur Lasermarkierung geeignete thermoplastische Harzzusammensetzung; und

(b) wenigstens ein Kupfersalz, ausgewählt aus der Gruppe bestehend aus Kupferphosphat, Kupfersulfat und
Kupferthiocyanat, wobei das Kupfersalz in einer Menge von 0,05 bis 10 Gewichtsteilen pro 100 Gewichtsteilen
der thermoplastischen Zusammensetzung vorhanden ist.

35 2. Thermoplastische Zusammensetzung nach Anspruch 1, dadurch gekennzeichnet, dass das Kupfersalz Kupfer-
phosphat oder Kupferthiocyanat ist.

40 3. Thermoplastische Zusammensetzung nach Anspruch 1, dadurch gekennzeichnet, dass das thermoplastische
Harz wenigstens ein Harz aufweist, ausgewählt aus Polycarbonat, Polyester, kautschukmodifizierten aromatischen
Monovinyldienharzen, Polyetherimide, Polyesteramiden, Polyamid, Polyestercarbonaten, Polyphenylensulfid, Po-
lyyamidimid, Polyetherestern, Polyetherimidestern, Polyarylat, Polymethylpenten, Polysulfonpolyethersulfon, Po-
lystyrol, kautschukmodifiziertes hochschlagzähes Polystyrol, Polyphenylenether, Polyetherketon, chloriertes Po-
lymer, fluoriertes Polymer, Flüssigkristallpolymer, Copolymeren der obigen oder Blends der obigen.

45 4. Thermoplastische Zusammensetzung nach Anspruch 4, dadurch gekennzeichnet, dass das thermoplastische
Harz umfasst Polyester, Polycarbonat, Polyestercarbonat, kautschukmodifizierte aromatische Monovinyldienhar-
ze, Polyetherimide, Polyesterester, Polyetherimidester oder Copolymeren oder Blends der obigen.

50 5. Thermoplastische Zusammensetzung nach Anspruch 5, dadurch gekennzeichnet, dass die thermoplastischen
Harze umfassen Polycarbonat,
Polyalkylenterephthalat, kautschukmodifizierte aromatische Monovinyldiene oder Copolymeren oder Blends der
obigen.

55 6. Thermoplastische Zusammensetzung nach Anspruch 6, dadurch gekennzeichnet, dass die Menge des Kupfer-
salzes sich zwischen 0,1 und 5 Gewichtsteilen der thermoplastischen Zusammensetzung bewegt.

7. Thermoplastische Zusammensetzung nach Anspruch 1, dadurch gekennzeichnet, dass die Menge des Kupfersalzes von 0,5 bis 3 Gewichtsteilen der thermoplastischen Zusammensetzung reicht.
- 5 8. Thermoplastische Zusammensetzung nach Anspruch 2, dadurch gekennzeichnet, dass die Partikelgröße des Kupfersalzes kleiner als 10 µm ist.
9. Thermoplastische Zusammensetzung nach Anspruch 9, dadurch gekennzeichnet, dass die Partikelgröße des Kupfersalzes kleiner als 1 µm ist.
- 10 10. Thermoplastische Zusammensetzung nach Anspruch 12, dadurch gekennzeichnet, dass die gesamte Farbabweichung zwischen der Basiszusammensetzung und des Teils, welcher das Laserlicht absorbiert, wenigstens 20 Einheiten ist, gemessen mittels Delta E.
- 15 11. Thermoplastische Zusammensetzung nach Anspruch 2, dadurch gekennzeichnet, dass die Zusammensetzung weiterhin Ruß oder Titandioxid aufweist.
12. Ein lasermarkierbarer Gegenstand, umfassend die Zusammensetzung gemäß Anspruch 1.
- 20 13. Verfahren zur Herstellung einer lasermarkierbaren thermoplastischen Zusammensetzung, bei welchem man die Zusammensetzung gemäß Anspruch 1 extrudiert.
14. Verfahren zur Herstellung eines lasermarkierbaren Gegenstands, bei welchem man:
- 25 (I) eine thermoplastische Zusammensetzung extrudiert und formt, umfassend:
- (a) wenigstens eine zur Lasermarkierung geeignete thermoplastische Harzzusammensetzung; und
- 30 (b) wenigstens ein Kupfersalz, ausgewählt aus der Gruppe bestehend aus Kupferphosphat, Kupfersulfat oder Kupferthiocyanat, wobei das Kupfersalz in einer Menge vorhanden ist die ausreicht, dass die lasermarkierbare thermoplastische Zusammensetzung Laserlicht außerhalb des sichtbaren Spektrums absorbiert, so dass der Teil der Zusammensetzung, welcher das Laserlicht absorbiert, eine sichtbar verschiedene und getrennt identifizierbare Farbe hat; und
- 35 (II) den Gegenstand mit einem Laser lasermarkiert, welcher eine Wellenlänge außerhalb des sichtbaren Spektrums hat.
15. Verfahren nach Anspruch 14, dadurch gekennzeichnet, dass die Wellenlänge des Lasers oberhalb 900 nm ist.

40 **Revendications**

1. Composition thermoplastique pouvant être marquée au laser, comprenant
- 45 (a) au moins une composition à base de résine thermoplastique appropriée pour un marquage au laser, et
(b) au moins un sel de cuivre choisi dans le groupe formé par le phosphate de cuivre, le sulfate de cuivre et le thiocyanate de cuivre, ledit sel de cuivre étant présent en une quantité comprise entre 0,05 et 10 parties en poids pour 100 parties en poids de composition thermoplastique.
- 50 2. Composition thermoplastique selon la revendication 1, dans laquelle ledit sel de cuivre est du phosphate de cuivre ou du thiocyanate de cuivre.
- 55 3. Composition thermoplastique selon la revendication 1, dans laquelle ladite résine thermoplastique comprend au moins une résine choisie parmi les polycarbonates, les polyesters, les résines monovinylidène-aromatiques modifiées par du caoutchouc, les poly(éther imide), les poly(ester amide), les polyamides, les polyester carbonates, les poly(phénylène sulfure), les poly(amide imide), les poly(éther ester), les poly(éther imide ester), les polyarylates, le polyméthylpentène, les polysulfones-poly(éther sulfone), le polystyrène, le polystyrène choc modifié par un caoutchouc, les poly(phénylène éther), les poly(éther cétone), les polymères chlorés, les polymères fluorés, les polymères de type cristaux liquides, ainsi que les copolymères et mélanges des polymères ci-dessus.

4. Composition thermoplastique selon la revendication 4, dans laquelle ladite résine thermoplastique comprend un polyester, un polycarbonate, un poly(ester carbonate), une résine monovinylidène-aromatique modifiée par du caoutchouc, un poly(éther imide), un poly(ester amide), un poly(éther ester), un poly(éther imide ester) ou un copolymère ou mélange des polymères ci-dessus.
5. Composition thermoplastique selon la revendication 4, dans laquelle ladite résine thermoplastique comprend du polycarbonate-poly(alkylène téréphthalate), une résine monovinylidène-aromatique modifiée par du caoutchouc ou un copolymère ou mélange de ces polymères.
10. 6. Composition thermoplastique selon la revendication 5 dans laquelle la quantité dudit sel de cuivre est comprise entre 0,1 et 5 parties en poids rapporté à la composition thermoplastique.
7. Composition thermoplastique selon la revendication 1, dans laquelle la quantité dudit sel de cuivre est comprise entre 0,5 et 3 parties en poids rapporté à la composition thermoplastique.
15. 8. Composition thermoplastique selon la revendication 2, dans laquelle la faille des particules dudit sel de cuivre est inférieure à 10 µm.
9. Composition thermoplastique selon la revendication 8, dans laquelle la taille des particules dudit sel de cuivre est inférieure à 1 µm.
20. 10. Composition thermoplastique selon la revendication 1, dans laquelle la différence de couleur totale entre la composition de base et la partie ayant absorbé la lumière laser est au moins égale à 20 unités de ΔE.
25. 11. Composition thermoplastique selon la revendication 1, dans laquelle ladite composition comprend en outre du noir de carbone ou du dioxyde de titane.
12. Article ouvré pouvant être marqué au laser comprenant la composition selon la revendication 1.
30. 13. Procédé de préparation d'une composition thermoplastique pouvant être marquée au laser comprenant l'extrusion d'une composition selon la revendication 1.
14. Procédé de préparation d'un article ouvré marqué au laser comprenant les étapes suivantes :
35. (I) extrusion et moulage d'une composition de résine thermoplastique comprenant
 - (a) au moins une composition à base de résine thermoplastique appropriée pour un marquage au laser, et
 - (b) au moins un sel de cuivre choisi dans le groupe formé par le phosphate de cuivre, le sulfate de cuivre et le thiocyanate de cuivre, ledit sel de cuivre étant présent en une quantité suffisante pour que ladite composition thermoplastique marquable au laser absorbe la lumière de laser ayant une longueur d'onde en dehors du spectre visible de manière à ce que ladite partie de ladite composition qui吸ue la lumière de laser a une couleur visible à l'œil et pouvant être identifiée séparément, et
40. (II) le marquage dudit article à l'aide d'un laser ayant une longueur d'onde en dehors du spectre visible.
45. 15. Procédé selon la revendication 14, dans lequel la longueur d'onde du laser est supérieure à 900 nm.

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